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22/30

### Regulation of Cellulase Gene Expression in the Filamentous Fungus Trichoderma reesei

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Basic features of regulation of expression of the genes encoding the cellulases of the filamentous fungus Trichoderma reasel QM9414, the genes chhi and chhi encoding cellobiohydrolasos and the genes egli, egli and egi5 encoding endoglucanases, were studied at the mRNA level. The cellulase genes were coordinately expressed under all conditions studied, with the steady-state mRNA levels of chil being the highest. Solka floc cellulose and the disaccharide sophorose induced expression to almost the same level. Moderate expression was observed when cellohiose or lactore was used as the carbon source. It was found that glycerul and sorbitol do not promote expression but, unlike glucose, do not inhibit it either, because the addition of 1 to 2 mM sophorose to glycerol or sorbitol cultures provokes high cellulase expression levels. These curbon sources thus provide a useful means to study cellulase regulation without significantly affecting the growth of the fungus. RNA slot blot experiments showed that no expression could be observed on glucose-containing medium and that high glucose levels abolish the inducing effect of sophorose. The results clearly show that distinct and clear-cut mechanisms of induction and glucose repression regulate cellulase expression in an actively growing fungus. However, derepression of cellulase expression occurs without apparent addition of an inducer once glucose has been depleted from the medium. This expression seems not to orise simply from starvation, since the lack of earbon or nitrogen as such is not sufficient to trigger significant expression.

A special feature of filamentous fungi and cellulolytic bacteria is the production of extracellular hydrolytic enzymes which provide them with the capability to use complex plant polysaccharides for growth. The major component of plant material, together with hemicellulose and lignin, is cellulose. It consists of long polymers of \$1,4-linked glucose units, which in turn form higher-order fibrillar structures. Cellulose is to a large extent in a crystalline water-insoluble form in plant material, and its hydrolysis consequently sets a challenge for the saprophytes. The cellulolytic organisms, bacteria and filamentous fungi, produce a set of enzymes which synergistically hydrolyze crystalline cellulose to smaller oligosaccharides and finally to glucose. Because cellulose is the most abundant organic carbon source on Earth, cellulolytic organisms play a very important role in carbon turnover in nature. The cellulolytic capability may also play a significant role in the ability of tungal plant pathogens or antagonistic fungi to attack their target organisms.

One of the most extensively studied cellulolytic organisms is the soft rot fungus Trichoderma reesei (reviewed in references 23, 41, and 56). Hypercellulolytic mutant strains secrete large amounts of cellulases, the largest published amounts being about 40 g/liter (9), and T. reesei therefore has also considerable industrial importance. The fungus produces a complete set of cellulases that are able to cleave the \$1,4-glycosidic bonds present in cellulose or cellulose derivatives. The cellulases are generally classified as cellobiohydrolases (CBH) (EC 3.2.1.91), which act as expenzymes and release cellobiose as a main product from crystalline cellulose; endoglucanases (EG) (EC 3.2.1.4), which have high affinity towards soluble cellulose derivatives and attack these by endoaction; and B-glucosidases (EC 3.2.1.21), which hydrolyse collooligosaccharides and the

The fungus produces high levels of cellulases on media containing cellulose or complex plant material, on cellubionolactone, and, to a variable extent, in the presence of the disaccharides cellobiose or lactose (reviewed in references 4 and 23). Production is not significant when glucose, fructose, or glycerol is used as the carbon source. A high level of cellulase expression is obtained when sophorose, a molecule consisting of two glucose units linked by a \$1,2-linkage, is added into the cultures (28, 37). This molecule could be formed from cellooligosaccharides by the transglycosylation activity of EGI (3, 8) or β-glucosidase (16, 58).

The fact that the major cellulase genes were cloned by differential hybridization showed that their expression is regulated at transcriptional level, and this has since been confirmed (1, 10-12, 31, 42). However, many hasic questions still remain to be thoroughly studied, such as whether the expression of the different cellulases is regulated by a shared mechanism and whether a low level of constitutive expression of cellulases occurs under all conditions. Constitutive expression might be needed for the initial release of a soluble inducing compound from the highly polymeric cellulose, which would further trigger high levels of cellulase expression. It has not been thoroughly studied whether mechanisms of true induction or repression are operating. The biochemical and molecular data available today is to some extent contradictory and has so far provided insufficient data to form the basis for further research directed toward, for instance, analysis of the functional regions in the promoters of the cellulase genes or the regulatory proteins involved. For these reasons, we have attempted to address some of the questions related to cellulase expression in filamentous fungi, and in this article we describe an investiga-

disaccharide cellohiose into glucose, providing the fungus with an easily utilizable carbon source for growth. Two genes encoding CBHs, cbh1 (50, 54) and cbh2 (5, 55), four encoding EGs, egil (38, 60), egil (46), egil (63), and egil (44, 45), and one encoding a \beta-glucosidase (2, 26) have now been reported from this organism.

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VOIL 63, 1997

CELLULASE EXPRESSION IN TRICHODERMA REESEI 1299

tion at the mRNA level of the basic mechanisms of cellulase regulation in the filamentous fungus T. reesei.

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#### MATERIALS AND METHODS

Strains and vectors. T. recrei QM9414 (29) was used throughout the studies. Plasmids pTTc1, pTTc9 (40), pTTc11 (39), pMS1 (46), pAS11 (44, 45) and pSV3 (61) were used as a source of the specific probes for the cbh1, cbh2, cgl1, cgl2, cgl5, and pgk1 genes, respectively. The T. recommend probe (30) probe was amplified from chromosomal DNA with sequence-specific primers.

Shake flask cultivations. The minimal media (MM) used consisted of 3% (lactose) or 2% carbon sources (glucose, cellobiose, sorbitol, or glycerol; when more than one carbon source was used, the concentration of each was 2%), 1.5% KH<sub>2</sub>PO<sub>4</sub>, 0.5% (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.06% MgSO<sub>4</sub>, 0.06% CnCl<sub>2</sub>, 0.0005% FcSO<sub>4</sub> · 7H<sub>2</sub>O. 0.00016% MnSO<sub>4</sub> · H<sub>2</sub>O. 0.00014% ZnSO<sub>4</sub> · 7H<sub>2</sub>O. and 0.00037% CoCl<sub>2</sub> · 6H<sub>2</sub>O. An organic altrogen source, 0.2% proteone peptone, was used in addition to aromonium when indicated. A total of 10° T. reseri CM0414 spores, collected from cultures grown on glucius and stored in 20% glycerol of -70°C, were inoculated into 50 ml of the medium in a 250-ml conical flask and grown at 28°C with shaking at 200 rpm. Four parallel flasks were inoculated for each medium and time point (14 to 92 h); the growth of the fungus, the p(t, and the sugar consumption were analyzed; and RNA was extracted.

Cultivation on cellulose-containing medium was done under the conditions described above. The fungus was grown for 49 h in minimal medium supplemented with 1% Solka fine cellulose and 0.2% peptone. At that time, glueree was added to 2%, and as a control, Solka fine cellulose was added to 3% final concentration, and the incubation was continued for 15 h before the mycelia were harvested. Additional control mycelia were grown for 72 h on 3% Solka fine cellulose and for 49 h on 1% Solka fine cellulose.

The cultivations in which supherose was used as an inducer were performed as follows. A total of 10<sup>7</sup> spores were inneulated into 50 ml of a culture medium containing either 5% gluense, 2% scribitol, or 2% glycerol and grown for 57 h (gluense) or 72 h (glycerol or scribital); I mM supherose was acted at that time and 10 h later, and mycelia were harvested 15 h after the first addition. To monitor the time course of chil mRNA accumulation, mycelia were harvested 3, 10, 24, and 48 h after addition of 1 mM sophorose to sorbital cultures.

In the experiment designed to study the effect of nitrogen or eathon starvation, the funges was grown in 250 ml of MM supplemented initially with 3% of glucuse in 2-liter shake flasks. At 48 h after inneulation, mycella from six parallel flasks were collected, washed, and transferred into three types of fresh medius medium similar to that used from the beginning, medium lacking the carbon source, and medium lacking the automorphism sulfate present normally in MM as the nitrogen source. The incubation was continued for a further 3 days. Samples for RNA isolation, determination of mycellal dry weight, pH, and ammonia and glucose amounts in the culture medium were taken 24, 36, and 48 h after the inoculation and 2, 5, 10, 24, 48, and 72 h after the transfer to the fresh media.

Bioreactor enlitvations. The bioreactor cultivation on glucose was performed in a medium initially containing 6% glucose, 0,1% yeast extract, 0.5% Bacto-Peptone, 0.4% KH2PC), 0.4% (NHa)2SO4, 0.05% MgSO4, 0.05% CxCl2. 0.0005% FcSO<sub>4</sub> - 7H<sub>2</sub>O, 0.00016% MnSO<sub>4</sub> · H<sub>2</sub>O, 0.00014% ZnSO<sub>4</sub> · 7H<sub>2</sub>O. and 0.00037% CoCl<sub>2</sub> · 6H<sub>2</sub>O in a 10-liter volume in a Chemap CF 2000 biogeactor at 29°C. Acreating was at 30 literalmin, and the pH was kept above 4.0 by automatic addition of NH<sub>2</sub>OH. The bloreactor was inoculated with mycella grown in glucose medium in shake flasks for 24 h in a 1-liter volume, and the glucose enneentration in the innershor was 0.8% at the time of inoculation. At 28 h of cultivation, additional yeart extract (5 g/fiter) and Bacto-Peptone (5 g/liter) were added. Also, 12% glucose was fed to the culture to keep the level of glucese above 2% between 30 and 50 h of cultivation. The total amount of giveose given during this period was 465 g. After that, the lungua was allowed to consume the remaining glucose from the medium. The total cultivation time was 144 h. The consumption of glucose and the growth of the fungus were manifered, and RNA was extracted at the time points indicated in Fig. 5.

The bloreactor cultivation on cellulose was performed in a 15-liter volume in a Champ CF 2000 bioreactor in the same medium as the gluence cultivation, except that gluence was replaced by 2% Solka floc cellulose at 29°C for 24 h and thereafter at 33°C. Acreation was at 30 liters/min. The initial pH was 5.2 without adjustment and was kept above 4.0 by automatic addition of NH<sub>4</sub>()11. The inoculum, 1 liter, was grown in the same medium for 24 h. The cultivation was continued for 67 h. and samples for RNA isolation were taken during growth at 20, 24, 29, 42, and 67 h.

Analysis of growth and consemption of sugars and ammonia. The growth of the fungus in the 250-ml shake flasks was measured by centrifuging the contents of one flask followed by lyaphilization overnight. From the higreactor cultivation on glucose medium and the 2-liter shake flasks, samples of 50 or 15 ml were filtered, washed, and baked at 100°C overnight.

The sugars left in the culture media were analyzed by high-pressure liquid chromatography (HPLC). The gluense concentration in the bioreactor cultivation was measured with the GOD-Perid test kit (Bochringer Mannheim). The ammonium concentration was measured with the ammonium test combination (Bochringer Mannheim).

Northern hint analyses. Total RNA was extracted from the mycelia as described previously (7). RNA was glyoxylated (47), run in a 1% agarose gel, and transferred to Hybond N sylon filters (Amendam). The hybridizations were performed in a hybridization solution containing 50% formanide, 1 M NaCl, 1% sudium clodecyl sulfate SDS, 10% decirate sulfate, 100 µg of herring sporm DNA per mi, and 2 to 5 × 10° cpm of the 32P-labelled DNA probes, which were either tail-length cDNA (chh1, cgb, and pgk1) or gene fragment (chh1, 660 hp; chh2, 660 hp; cgl1, 660 hp; cgl2, 820 hp; and acc, 680 hp) probes. The RNA amount toucked on the gel was controlled by scriding orange staining. Hybridization signals (see Fig. 2 and 3) were quantified with a Phosphorimager (Molecular Dynamics), and cellulase signals were numalized to actin levels for comparisons.

Slot blot analysis of RNA. Total RNAs were diluted into appropriate extent-trations with water and glyonylated by adding the same volume of the glyonal mixture (64). After incubation for 1 h at 50°C, a dilution series was made and the samples were applied to a Hybond N nylon filter by using the Bio-Dot SF Electing apparatus (Bin-Rail) as specified by the manufacturer. Filters were hybridized as described for the Northern blot analyses. The probe fragments were amplified by PCR with specific primers to obtain fragments of equal length from each gone, and these were labelled to a similar specific activity (3 × 10° cpm/µg of DNA).

#### **RESULTS**

Induction of cellulase expression. The media usually used to promote high cellulase expression are based on mixtures of plant materials or cellulose. These materials interfere with the estimation of dry weight and isolation of RNA and are also often undefined in nature. For these reasons, the first shake flask experiments were carried out on medium containing 2% cellobiose as the only carbon source, and the results were compared to those obtained with 2% glucose cultures. Mycelial dry weight, pH, carbon source utilization, and expression of the glycolytic phosphoglycerate kinase gene, pgk1, as a general control of metabolism (62) were analyzed regularly from the cultures.

Cellobiose clearly induced the expression of chil (Fig. 1A) and, in a similar manner, that of the other cellulase genes (data not shown) after a rather long lag period. Reduced expression was observed at a later stage of growth, when glucose was accumulating although cellobiose was still present. No cellulase transcripts were detected in similar cultures on glucosecontaining media (Fig. 1C). The presence of an organic nitrogen source has been suggested to be important for cellulase production on Solka floc cellulose medium (25), and the effect of addition of peptone to cellobiose cultures was analyzed. This did not improve expression (Fig. 1B); on the contrary, expression was barely detectable. This result indicates that peptone seems not to enhance induction per se, and the earlier reports on its positive effect on cellulase production could instead be due to increased growth of the fungus and subsequent elevated cellulase levels in production conditions.

It has been suggested previously (see Discussion) that collulase expression would occur on carbon sources supporting only poor growth. The expression patterns observed in the collobiose cultures would be in accordance with this: lower expression was obtained in the collobiose-peptone culture, where growth was clearly more rapid than on the collobiose medium in which ammonium was the only nitrogen source. However, no expression occurred in a similar shake flask experiment carried out on 2% sorbitol medium (Fig. 2, lanes 2 and 3; Fig. 3, lane 1), where growth was also poor, even poorer than on collobiose (data not shown).

Additional studies showed that scribitol may be used as a nonrepressing, noninducing carbon source. As mentioned above, no expression occurred on sorbitol medium but addition of sophorose to the culture medium efficiently induced cellulase expression (Fig. 2, lanes 2 to 4). Sophorose (1 mM) was added twice to the sorbitol culture, at 72 and 82 h, and mRNA analyses were performed 15 h after the first addition, when over 1 mM sophorose was still present in the medium.

1300 ILMÉN ET AL.

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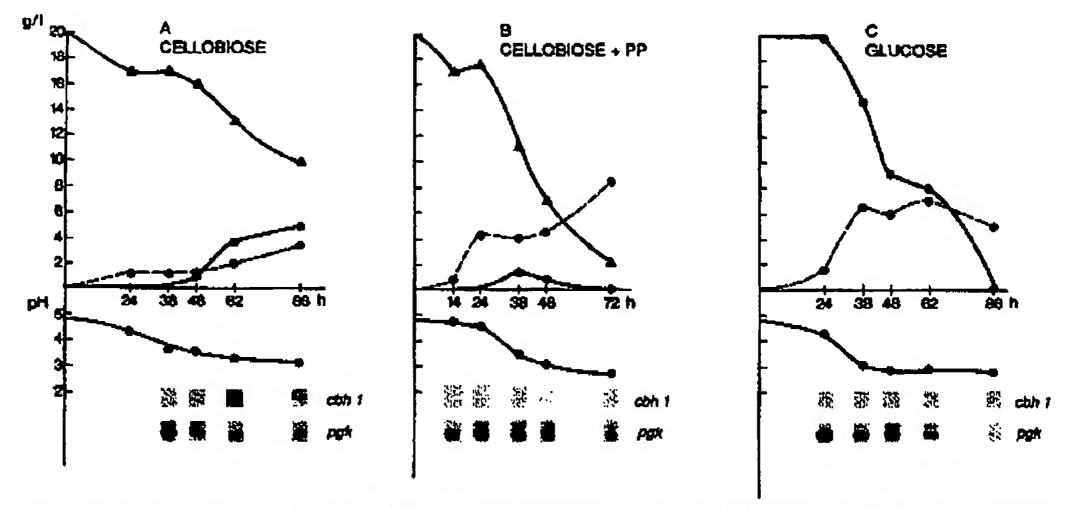


FIG. 1. Shake flask cultivations on media containing 2% celloblese (A). 2% celloblese plus protesse persone (PP) (B), and 2% glucose (C). The amounts of celloblese (A) and glucose (——), the fungal dry weight (——), and pH of the medium (———) were analyzed and samples for RNA isolations were taken at the time points indicated. Northern analyses of total RNA (3 µg) using ch11 and pok1 probes is shown below.

Strong sophorose induction of the cellulases chh1 and chh2 was observed, and egi1, egi2, and egi5 were also induced, although less efficiently. No mRNAs could be detected at different stages of growth without addition of the inducer (Fig. 2, lanes 2 and 3). Also, the presence of 2% cellobiose in sorbital cultures from the beginning of the cultivation provoked some cellulase transcription (Fig. 2, lane 1), and this level was comparable (data not shown) to that obtained in the cultivations carried out on cellobiose alone (Fig. 1).

That prolonged strong sophorose induction of cellulases can be obtained by adding small amounts of sophorose to sorbitol cultures under the above-mentioned conditions was further shown in a separate set of experiments in which the time course of sophorose induction was investigated (Fig. 3). Moderate cbh1, cbh2, egl1, and egl5 expression was detected 3 h after the addition of sophorose, and the mRNAs continued to accumulate and remained abundant until at least 48 h. Sophorose induction of egl2 was observed in a longer exposure. These steady-state mRNA levels are comparable to those obtained when Solka floc cellulose is used as the sole carbon source as shown in Fig. 2 (lanes 4 and 10 to 12).

The finding that sorbitol can in practice he considered a neutral carbon source for cellulase expression prompted us to study the nature of glycerol in this respect. Sophorose was added twice to 2% glycerol cultures in a similar manner to that described above for the sorbitol cultures. A high level of cellulase expression occurred; the data for the five cellulases are shown in Fig. 2, lanes 5 to 9. Sophorose induction of cellulases was observed in cultures containing 5% sorbitol or glycerol as well (data not shown). Thus, both sorbitol and glycerol can be considered "neutral" carbon sources with respect to cellulase expression in contrast to glucose (see below).

The comparison between the RNA samples collected from the different growth conditions presented in Fig. 2 revealed that the highest expression levels of all the cellulase genes studied were reached in glycerol-sophorose cultures, closely followed by Solka floc cellulose and sorbitol-sophorose (Fig. 2. lanes 4 and 8 to 12). The differences between these highly induced conditions were two- to fourfold. In the sorbitol-cellubiose cultures (Fig. 2, lane 1), the expression levels were about 30-fold lower than under the highly induced conditions.

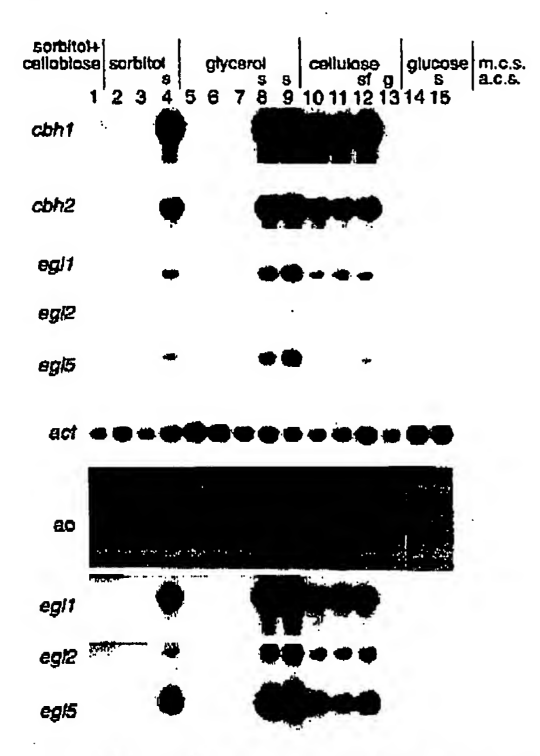
Relative expression levels of the cellulases. To study in more detail the relative transcript amounts of the cellulases cbhl, chh2, and egl1, which are considered the three major cellulases, RNAs from mycelia grown under different inducing conditions were compared by a slot blot analysis with probes of the same length and specific activity. Figure 4 shows the results from the shake flask cultivations carried out on sorbitol-sophorose medium and on 3% lactose medium. Also, samples from a bioreactor cultivation on Solka floc cellulose containing medium were included in the analysis. The steady-state mRNA levels of chhl were the highest of the cellulases under all conditions, followed by chh2 and cgl]. chh] mRNA was approximately 1.5 and 3 times more abundant than chh2 and egil mRNAs. respectively, as determined by laser scanning densitometry, although the visually estimated differences seemed larger (Fig. 4). The mRNA levels on lactose medium remained clearly lower than on sorbitol-sophorose and cellulose media, with the difference in steady-state mRNA levels between lactose and cellulose being about 60-fold. In a separate slot blot experiment, the expression level obtained on the 2% cellobiose medium was shown to be comparable to that of the lactose culture (data not shown). This is well in accordance with the 20-fold difference reported at the protein level between cellulose and cellabiase cultures (27) and the estimations presented above based on results in Fig. 2. It is noteworthy that as observed in the shake flask experiments (Fig. 2), the induction levels after 15 h of sophorose induction approach those obtained on Solka floc cellulose medium after 20 h in a bioreactor.

In the bioreactor cultivation on Soika floc cellulose medium, the cellulase transcript levels were clearly higher than the mRNA levels of the phosphoglycerate kinase gene, pgk1, in-

1301

CELLULASE EXPRESSION IN TRICHODERMA REESEI

Vot., 63, 1997



650-845-6504

FIG. 2. Induction of chh1, chh2, chl2, chl2, and ch45 expression by sophorose and collabiose in cultures containing surfittel and glycerul and repression in the presence of glucose. RNA was isolated from mycelia grawn an media containing 2% norbitol plus 2% cellobione (lane 1). 2% norbitol (lanes 2 to 4). 2% glycerol (lanes 3 to 9), Solks flux cellulose (lanes 10 to 13), or 5% glucuse as the main carbon source (m.c.s.), Additional carbon sources (a.c.s.) were added to relected cultures as follows: I mM sophorese (s) was added to antiltol and glycerol cultures at 72 and 82 h and to glucose cultures at 57 and 67 h. Mycelia were collected 15 h after the first addition. Glucose (g) was added to 2% and Solka fine cellulose (sf) was added to 3% (ma) concuntration at 49 h of the cellulose-based cultivation, and the mycelia were collected 15 h later. A 5-µg portion of total RNA was loaded onto the gel. Four identical blots were prepared and hybridized simultaneously with the cellulase probes, chh1, chh2, and cg11 probes were of the same length and specific activity (1 × 10° cpm/µg); cg/2, and cg/5 probes were less serive  $(2 \times 10^6 \text{ cpm/µg})$ . Control hybridizations with an actin probe (act) were done after stripping the membranes, and the result is shown for one of the gels together with the scridine orange staining (ao). The filters were exposed for the same time (four top panels). Longer exposures of the endogluesnase hybridizations are shown in the two lowest panels. Lanes: 1, sorbited plus cellubiuss, 72 h; 2. sorbitol, 72 h; 3, sorbitol, 87 h; 4, sorbitol plus sophorose, 87 h; 5, glycerol, 72 h; 6, glycerel, 87 h; 7, glyceral, 92 h; 8, glycerol plus sopharase 87 h; 9, glycerol plus sophorose, 92 h; 10, 3% Solka floe cellulose, 72 h; 11, 1% Solka floe cellulose, 49 h; 12, 1% plus 2% Solka floc cellulose, 64 h; 13, 1% Solka floc cellulose plus 2% glucose: 14, 5% glucose, 72 h; 15, glucose plus sophiruse.

volved in glycolysis, with the difference between chh1 and pgk1 heing roughly 50-fold. On the other hand, on cellobiose medium, where induction of cellulases is weaker, the steady-state expression level of pgk1 exceeded that of the cellulases (data not shown).

Gincose repression of cellulase expression. Data from the shake flask cultivations showed that glucose acts as a repressor of cellulase synthesis at the transcriptional level. No transcrip-

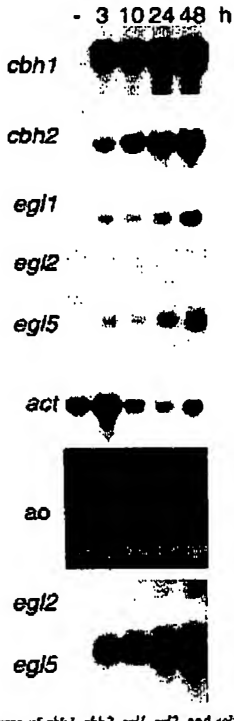
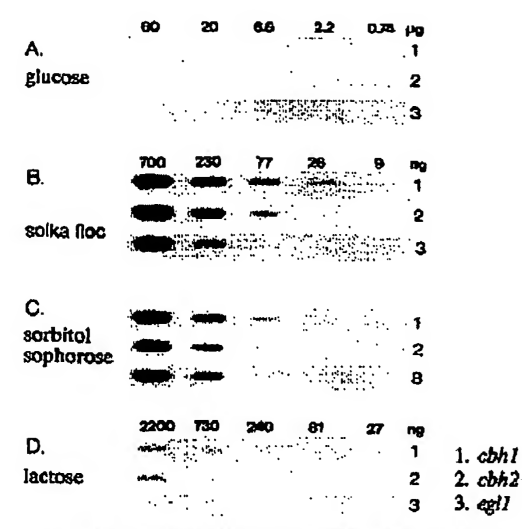


FIG. 3. Time course of chil, chh2, egil, egil, and egil mRNA accumulation after suphorose addition to 2% surhitol cultures. Incubation was carried out for 72 h on surhitol; then 1 mM suphorose was added, and RNA was isolated at the time points indicated after sophorose addition (3, 10, 24, and 48 h) and before suphorose addition (-). Northern blot analysis was performed exactly as described in the legend to Fig. 2 and hybridized with the probes indicated on the left. The filters were exposed for the same time (four top panels). A longer exposure is included to demonstrate the egil signal (lowest panel).

tion was observed if sophorose was added to a glucose-based culture (Fig. 2, lanes 14 and 15). The repressing action of glucose on cellulase expression was further shown by addition of glucose to fully induced cellulose-based cultures (lanes 10 to 13), which resulted in the disappearance of the cellulase transcripts.

To address further the question of possible low basal levels of expression on pure glucose, T. reesei QM9414 was grown in a bioreactor (Fig. 5). This allows the glucose level to be carefully controlled and several RNA samples to be analyzed at different time points of the culture. No signals with the cellulase probes cbh1, cbh2, and egl1 were detected in Northern blot analyses at different time points when glucose was present in the culture (Fig. 6, time points 35 to 75 h). However, the presence of low constitutive expression levels cannot be necessarily excluded on the basis of a regular Northern blot analysis, and for this reason a sample from the 35-h time point was included in the same dot blot analysis as described above (Fig. 4). No cellulase transcripts were detectable, even when overloading the analysis with RNA. Comparison of the different culture conditions shows that the steady-state cellulase transcripts would be at least 1,000-fold (in the case of cbh1, roughly 1302 ILMÉN ET AL

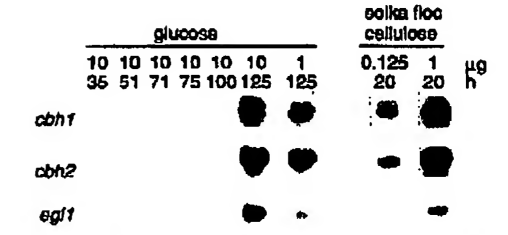




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FIG. 4. Slot blot analysis on steady-state cellulase transcript levels produced under various culture cunditions. Total RNA (0.74 to 60 µg) from glucose-grown mycelia (A) (hioroactur cultivation, 35 h), total RNA (9 to 700 ng) from Solka floc cellulase-grown (B) (hieroactor cultivation, 20 h) and sorbitol-sophorose grown (C) (shake flost cultivation, 15 h after sophorose addition; total time, 87 h), mycelia, and total RNA (27 to 2200 ng) from lactose-grown mycelia (D) (shake flost cultivation, 96 h) were loaded. Cellulase probes of equal length were labelled to the same specific activities and are indicated on the right by numbers: L chiri; 2, chir2, 3, cg/f.

6,000-fold) more abundant in the bioreactor cultivation on Solka floc cellulose medium than in the bioreactor cultivation on glucose, provided that expression occurred on glucose but was not detectable by the method applied. Addition of an inducer did not change the situation, since no mRNA was



PIG. 6. Northern blot analyses of expression of the cellulose genes chil, chi2, and call during bioreactor cultivation of glucose containing medium at different time points. For comparison, samples from a Solka floc cellulose cultivation in a bioreactor (20 h) are shown on the right. Total RNA at 0.125 to 10 µg was loaded onto the gel as indicated above each lane.

detected when 60 µg of RNA from the previously described 5% glucose-sophorose shake flask cultivation (Fig. 2, lane 15) was analyzed in a similar dot blot analysis (data not shown). These data show that glucose repression of cellulase expression is strong in *T. reesei* QM9414. No cellulase mRNAs were observed when RNAs were analyzed similarly from the fungus grown on sorbitol medium (data not shown).

Analyses of multiple RNA samples from glucose-grown mycelia have always produced negative results with cbh1, cbh2, cg/1, and cg/2 probes. This has usually been the case with cg/5 hybridizations, except that in a few cases a positive signal has been detected (data not shown), and the significance of this observation should be studied further.

The fungus was allowed to consume the glucose completely in the bioreactor cultivation described above (Fig. 5), and RNA samples were also analyzed from this period. Surprisingly, after a lag period of roughly 45 h after glucose depletion from the culture, relatively high levels of chh1, chh2, and egil mRNAs were detected in the Northern analyses (Fig. 6). The mRNAs were already slightly visible at the 100-h time point, although they were not detectable with the exposure time shown in Fig. 6. At the 125-h time point, the expression levels

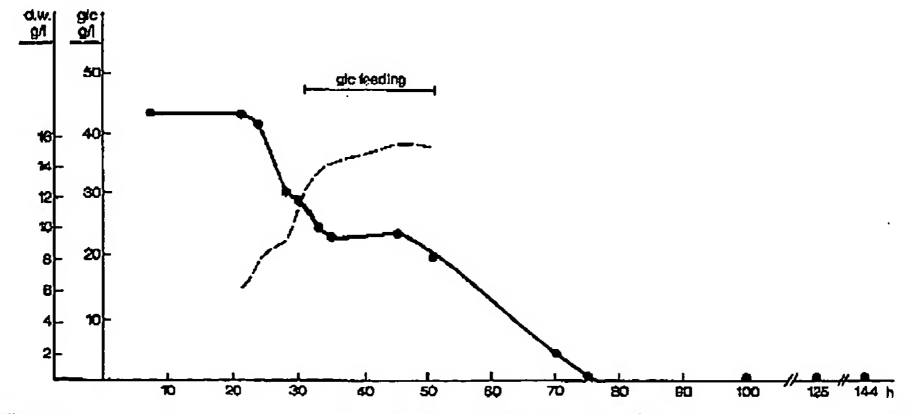


FIG. 5. Bistremeter cultivation on glucine-extrationing medium. The glucose concentration in the medium (solid line) and the dry weight of fungus (dashed line) were measured, and RNA was isolated at the time paints indicated by dots. Concentrated glucose solution was fed into the hierescipe between 30 and 50 h of cultivation, after which the fungus was allowed to consume all the glucose from the medium.

CBILLULASB EXPRESSION IN TRICHODERMA REESEI

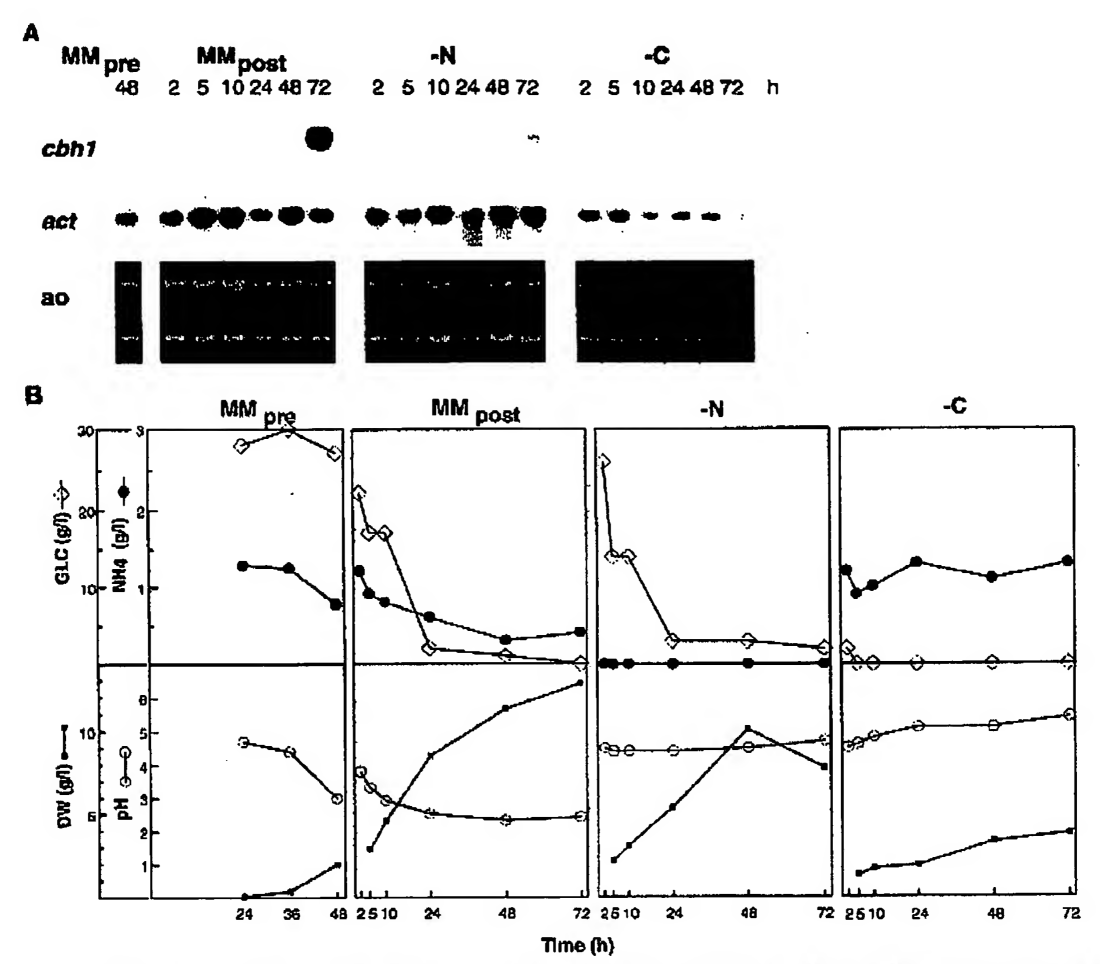


FIG. 7. Northern analysis on chil expression (A) and course of the shake flask cultivations (B) on minimal medium and on media lacking either a nitrogen or a carbon source. The fungus was grown for 48 h in minimal medium containing 30 g of glucose per liter and 1.3 g of ammonia per liter (MMpre). Thereafter, the mycelia were harverted, washed, and divided into fresh minimal medium supplemented with both glucose and ammonia (MM<sub>post</sub>) and into minimal medium lacking nitrogen (¬N) or earlien (¬C), and the incubation was continued for a further 72 h. Amounts of glucose (GLC) and ammonia, p11 of the medium, and mytelial dry weight (DW) were determined, and RNA was isolated at the time points indicated. A 2-µg portion of RNA was loaded outo the gel. Hybridization to an actio probe (ect) is shown as a control, an, acriding orange-stained gel,

were roughly 10% of those detected in the bioreactor cultivation on the cellulose medium shown for comparison. This result demonstrates that cellulase expression can be obtained after glucose depletion without an apparent addition of an inducing compound. We had earlier observed that this kind of derepression also occurs in prolonged shake flask cultivations on 2% glucose medium, especially if peptone is included, occurring then after about 24 h of glucose depletion (data not shown).

To further study if carbon or, alternatively, nitrogen depletion as such is provoking cellulase expression, the fungus was grown in shake flasks in a normal minimal medium supple-

mented with glucose and containing ammonium sulfate as a nitrogen source. In an active growth phase, the mycelia were washed and transferred into three different media, one the same as that used from the beginning (to serve as a control), one depleted of carbon, and one depleted of nitrogen (Fig. 7). Expression of chh1 was not detected when analyzed at different time points between 2 h and 2 days after the transfer. Only 3 days after the transfer, cb/1 mRNA was produced in the control cultures containing glucose and ammonia or in the culture containing glucose but depleted of nitrogen (Fig. 7A). This coincided with very low glucose levels in the medium. Expression of chal was not detected in the culture transferred to a

1304 ILMÉN ET AL. APPL ENVIRON MICROBIOL

medium lacking the carbon source (Fig. 7A), although, for instance, the starvation-inducible hydrophobin mRNA has already appeared after 2 to 10 h in the same experiment (34). The results obtained with cbh2 and egl1 probes (data not shown) were identical to cbh1. The RNA yield was very low at the late stages of the experiment in the medium without a carbon source (Fig. 7B), indicating that there were only few viable cells, which made it impossible to continue the experiment any further. These results suggest that carbon starvation per se is not sufficient to initiate cellulase expression, because complete removal of the carbon source from the medium had no effect even after 3 days of incubation.

#### **DISCUSSION**

Collulose and sophorose exerted the highest levels of collulase transcription in T. reesei QM9414, followed by moderate levels on lactose and cellobiose and no expression on glucose, glyccrol, or sorbitol. The results give a general picture of the potential of cellulase expression on these different carbon sources and are in accordance with collulase protein levels generally obtained. The relative transcript levels of the different cellulase genes with that of cbh I being the highest, are also comparable to the amounts of the specific proteins produced in the culture medium (16). It was suggested previously (19, 27) that collulase expression would not occur on carbon sources that promoted rapid growth, such as glucose and glycerol, but would be provoked on poor carbon sources, such as cellulose, cellobiose, and lactose. Our results do not show correlations between growth and cellulase expression, a conclusion also reached by others (31). Rather, our results show that distinct mechanisms of induction and repression are operating in the cellulase expression in Trichoderma.

The results clearly show the strong inducing power of sophorose when added at 1 to 2 mM to the culture; this was studied at the mRNA level previously (10, 12). Sophorose is consequently a good candidate for the natural inducing compound for cellulase expression. The reports on the use of cellobiose as a carbon source for cellulase expression are, on the other hand, to some extent controversial. Cellobiose has been considered a poor inducer (13, 15, 16, 37, 43) but is also to exert as good induction as cellulose if growth is kept slow and the level of cellobiase is kept low by controlled feeding of collobiose into the culture (59). In the cultivations presented here, collobiose promoted cellulase transcription to a moderate level comparable to that of lactose, which shows the inducing effect of cellobiase or al sophorose formed from it. The difficulty with this carbon source is, however, evident, as induction is not immediate and seems to be variable depending on the culture conditions. The failure to provoke cellulase expression could be explained, on one hand, by the lack of an inducing effect when consumed rapidly from the culture, as observed in this study in the presence of peptone, or, on the other hand, by the accumulation of glucose and subsequent glucose repression of collulase transcription. The inducing power could thus be dependent on the ratio between sophorose and glucose formation by  $\beta$ -glucosidase and their uptake from the medium. It is also likely that organic nitrogen sources such as peptone are used as both nitrogen and carbon sources (see also reference 21) and could somehow interfere with cellulase expres-

As reported previously (10), no cellulase mRNAs could be detected in fungus grown on glycerol as the only carbon source. However, glycerol has not been previously studied at the molecular level in the presence of an inducer, and the literature available groups this carbon source with glucose as being a

compound inhibiting cellulase expression (see reference 21 for discussion). Our results clearly show that glycerol, as well as sorbitol, does not as such promote expression but does not inhibit it either, because addition of sophorose to glycerol or sorbitol cultures provokes high cellulase transcript levels. Thus, the cellular effects exerted by glucose and glycerol-sorbitol are clearly distinct. This is in agreement with the situation in the yeast Saccharomyces cerevisiae (14). Studies on cellulase regulation are very much hampered by the fact that the most natural carbon sources promoting cellulase expression, such as cellulose, are also used for growth by the fungus. The use of glycerol and sorbitol as carbon sources now allows powerful and rapid cellulase gene induction to be obtained in growing cultures with 1 to 2 mM sophorose without significantly affecting the growth of the fungus.

It is generally believed that small soluble molecules are released from cellulose and act as inducing compounds or are converted to such. To explain this initial attack, cellulase expression has been suggested to occur constitutively at low levels under all conditions including only glucose and no inducer as a carbon source, and evidence for this has been presented (10, 20, 22, 31). The analysis carried out in this work shows that glucose repression is tight, and the differences in mRNA levels between fully induced and repressed states are great in an actively growing fungus provided that the presence of glucose in the culture is carefully controlled. It is still possible, however, that biologically relevant mRNA levels remain undetected in experiments based on steady-state mRNA levels if the mRNA turnover on glucose-based media is especially rapid. Mechanisms of glucose repression are in any case operating in the regulation of cellulase expression. We have recently shown by site-specific mutagenesis of the cbh1 promoter that regions similar to the binding sites of the regulatory proteins MIG1 of S. cerevisiae (35) and CREA of Aspergillus nidulans (24), involved in glucose repression in these organisms, seem to be at least partially responsible in vivo for the glucose repression of cbh1 (18). The Trichoderma equivalent of creA, cre1, has recently been cloned (17, 52, 53). The demonstration that cellulase expression occurs on glucose-containing medium in the hypercellulolytic T. reesei strain Rut-C30 and that this is caused by a mutant form of the crel gene (17) further confirms glucose repression of cellulase expression. The earlier reports pointing toward constitutive expression on pure glucose could be explained by the use of the mutant strain Rut-C30 in the studies or, alternatively, by the release of glucose repression once glucose has been consumed from the medium. This kind of derepression was clearly seen in both shake flask and bioreactor cultivations carried out in this work.

This capability of cellulase expression after glucose depletion, without the addition of an inducer, could provide the fungus with a survival mechanism under starvation conditions. Starvation has previously been suggested to induce cellulase synthesis (15). Interestingly, however, depletion of glucose as such was clearly not enough to initiate cellulase expression, as demonstrated by a prolonged incubation of pregrown washed mycelium in a medium containing no carbon source. This suggests that some kind of induction is required, and this might be related to growth since cellulase expression occurred in cultures where growth had occurred (Fig. 7). It is possible that oligosaccharide molecules released from the cell walls of the starving fungus act as inducing compounds or are converted to them. It is also possible that sophorose or some other inducing sugar is formed from the glucose present in the culture by the transglycosylation activity of the possible constitutive β-glucosidase (51, 57) and will later exert induction after glucose has heen depleted and repression released. The latter alternative is

1305

Vol. 63, 1997

supported by the fact that cellulase expression was not deteeted in mycelia incubated without a carbon source but was detected in glucose-grown mycelia (starved or not starved for nitrogen) after glucose was depleted from the medium. However, no sophorose could be detected in the medium in the glucose-based bioreactor cultivation by HPLC, which indicates that if extracellular sophorose was present, its level was below

the detection limit, 30 µM. Conidium-hound cellulases (22, 32) could provide another means for the fungus to attack cellulose in the absence of an inducer. No indication for conidiation was, however, observed at the later stages of the cultivations prosented in this work.

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Earlier mutant data suggest that separate regulation could exist for B-glucosidases, cellobiohydrolases, and endoghicanases (9, 33, 36, 48, 49). Our data show that the major cellulases, cellobiohydrolases I and II and the endoglucanases I and II of T. reesei, are coordinately expressed under all the conditions studied. This also seems to be the case in the derepressed stage after glucose depletion. These results indicate a common regulatory mechanism which retains the proportions of these cellulases. This, however, does not contradict the possibility that specific regulatory mechanisms are revealed in a mutant situation which, in the wild-type situation, all contribute to the overall control, keeping it apparently similar for all cellulases under standard conditions. Furthermore, T. reesei still produces at least one other \(\beta\)-glucosidase (6) and an additional endoglucanase, EGIII (63). Consequently, it is still possible that these cellulases, EGV, or some yet unknown enzymes, escape common regulation and are more constitutively expressed, thus having the potential of immediate release of an inducer once cellulose is present.

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1306 ILMÉN ET AL.

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